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SETOGUCHI NOBURO**(54) **CONDENSED IMIDAZOLE DERIVATIVE**

(57) Abstract:

NEW MATERIAL: A compound expressed by formula I (ring A is benzopyran ring, tetrahydroquinoline ring or indoline ring; which may have a substituent group; R is phenyl or pyridyl which may have a substituent group; n is 0 or 1 or an acid addition salt thereof.

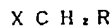
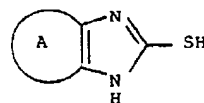
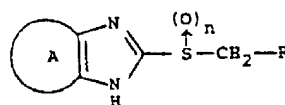
EXAMPLE:

2-[(3-Methyl-2-pyridyl)methylthio]pyrano[3,2-e]benzimidazole.

USE: An antiulcer agent, analgesic agent and anti-inflammatory agent.

PREPARATION: A compound expressed by formula II is reacted with a compound expressed by formula III (X is halogen) in the presence of a base, e.g. NaHCO_3 , NaOH or pyridine, in a solvent, e.g. benzene, methanol, DMF or water, at 60°C to 110°C for 20 min to 6 hr to afford the aimed compound expressed by formula I etc.

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